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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/252,828	02/19/1999	KE-WEN DONG	024754/0114	4940
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WILMER CUTLER PICKERING HALE AND DORR LLP THE WILLARD OFFICE BUILDING 1455 PENNSYLVANIA AVE, NW			VENCI, DAVID J	
			ART UNIT	PAPER NUMBER
WASHING	TON, DC 20004		1641	
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Please find below and/or attached an Office communication concerning this application or proceeding.

	Application No.	Applicant				
	Application No.	Applicant(s)				
Office Action Summary	09/252,828	DONG ET AL.				
Onice Action Summary	Examiner	Art Unit				
The MAILING DATE - SAL'-	David J Venci	1641				
The MAILING DATE of this communicate Period for Reply	ion appears on the cover sheet wi	th the correspondence address				
A SHORTENED STATUTORY PERIOD FOR THE MAILING DATE OF THIS COMMUNICA  - Extensions of time may be available under the provisions of 3 after SIX (6) MONTHS from the mailing date of this communic  - If the period for reply specified above is less than thirty (30) da  - If NO period for reply is specified above, the maximum statuto  - Failure to reply within the set or extended period for reply will, Any reply received by the Office later than three months after the earned patent term adjustment. See 37 CFR 1.704(b).	TION. 7 CFR 1.136(a). In no event, however, may a ration. ys, a reply within the statutory minimum of third y period will apply and will expire SIX (6) MON by statute. cause the application to become AE	eply be timely filed  by (30) days will be considered timely.  ITHS from the mailing date of this communication.  SANDONED (35 U.S.C. § 133)				
Status						
1) Responsive to communication(s) filed o	n <u>18 May 2004</u> .					
2a)⊠ This action is <b>FINAL</b> . 2b)[						
3) Since this application is in condition for	·					
closed in accordance with the practice u						
Disposition of Claims						
4) Claim(s) 69-78 is/are pending in the app	olication.					
	4a) Of the above claim(s) <u>73-78</u> is/are withdrawn from consideration.					
5) Claim(s) is/are allowed.						
6)⊠ Claim(s) <u>69-72</u> is/are rejected.						
7) Claim(s) is/are objected to.						
8) Claim(s) are subject to restriction	and/or election requirement.					
Application Papers						
9)⊠ The specification is objected to by the Ex	kaminer.					
10)⊠ The drawing(s) filed on <u>03 October 2003</u>		ccepted or b) objected to by the				
Examiner.						
Applicant may not request that any objection	ı to the drawing(s) be held in abeyan	ce. See 37 CFR 1.85(a).				
Replacement drawing sheet(s) including the						
11)☐ The oath or declaration is objected to by						
Priority under 35 U.S.C. § 119						
12) Acknowledgment is made of a claim for the	oreign priority under 35 U.S.C. §	119(a)-(d) or (f).				
a) ☐ All b) ☐ Some * c) ☐ None of:						
1. Certified copies of the priority doc	uments have been received.					
2. Certified copies of the priority doc	uments have been received in A	pplication No				
3. Copies of the certified copies of the	ne priority documents have been	received in this National Stage				
application from the International		J				
* See the attached detailed Office action fo	r a list of the certified copies not	received.				
Attachment(s)						
<ol> <li>Notice of References Cited (PTO-892)</li> <li>Notice of Draftsperson's Patent Drawing Review (PTO-90)</li> </ol>	ummary (PTO-413)					
2) Notice of Draftsperson's Patent Drawing Review (PTO-948)  3) Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08)  Paper No(s)/Mail Date  Notice of Informal Patent Application (PTO-152)						
Paper No(s)/Mail Date .	6) Other:	A Control of the Cont				

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#### **DETAILED ACTION**

Examiner acknowledges Applicants' amendment filed 11/19/03, which canceled all pending claims (i.e. claims 48-68) and added new claims 69-78.

Examiner acknowledges Applicants' supplemental amendment filed 5/18/04, which made several amendments to the specification, sequence listing, claims, and drawings.

Currently, claims 69-78 are before the Office.

### Election/Restrictions

Newly submitted claims 73-78 are directed to an invention that is independent or distinct from the invention originally claimed.

Claims 73 and 74 are directed to polypeptides and glycopolypeptides comprising materially different amino acid sequences as evidence by separate amino acid position substitutions. These separate polypeptides and glycopolypeptides bear distinct structural or biochemical properties as substantiated by the separate amino acid substitution positions thereby having different binding characteristic and functionality. Therefore, each disclosed patentably distinct glycopolypeptide is considered a separate invention and changes the scope of the previously elected invention.

Claims 75-78 are directed to a human ovarian cell containing a vector.

Since applicant has received an action on the merits for the originally presented invention, this invention has been constructively elected by original presentation for prosecution on the merits. Accordingly, claims 73-78 are withdrawn from consideration as being directed to a non-elected invention. See 37 CFR 1.142(b) and MPEP § 821.03.

## Specification

The disclosure is objected to because of the following informalities:

The substitute raw sequence listings filed November 14, 2000 and May 18, 2004 are objected to under 35 U.S.C. 132 because they introduce new matter into the disclosure. Under 35 U.S.C. 132, no amendment shall introduce new matter into the disclosure of the invention. The added material that is not supported by the original disclosure can be found under the listing for SEQ ID NO: 3. Specifically, the description of the organism has been changed from "mouse" to "murine sp." Applicant is required to cancel the new matter in the reply to this Office Action.

In addition, Applicants' have amended the specification to include reference to a "Table 1" (p. 15, line 2). See also p. 10, line 13. Currently, there is no such table of record. If applicants wish to add a table to the specification at a later date, applicants must pay careful attention not to add new matter. Clarification is required.

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Claim Rejections - 35 USC § 101

35 U.S.C. 101 reads as follows:

Whoever invents or discovers any new and useful process, machine, manufacture, or composition of matter, or any new and useful improvement thereof, may obtain a patent therefor, subject to the conditions and requirements of this title.

Claims 70 and 72 are directed to non-statutory subject matter. Specifically, claims 70 and 72 read on products of nature. Amendment of these claims to recite "isolated" or "recombinantly produced" polypeptides or glycopolypeptides will obviate this rejection.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

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Claims 69-72 are rejected under 35 U.S.C. 103(a) as being unpatentable over Chamberlin & Dean, 87 PROC. NATL. ACAD. Sci. 6014 (1990) in view of Kinloch et al., 92

PROC. NATL. ACAD. Sci. USA 263 (1995).

Chamberlin & Dean teach a human ZP3 protein comprising the amino acid sequence of

the claimed SEQ ID NO: 2 (See p. 6017, Fig. 3).

Chamberlin & Dean do not teach a recombinantly produced polypeptide. Chamberlin &

Dean also do not teach a glycopolypeptide.

However, Kinloch et al. teach a recombinantly produced ZP3 polypeptide and

glycoprotein (See p. 264, GLYCOPROTEIN PURIFICATION) in order to create and test

mutant ZP3 proteins for sperm-binding activity.

Therefore, it would have been obvious for a person of ordinary skill in the art to combine

the ZP3 amino acid sequence of Chamberlin & Dean with the method of making

recombinant ZP3 protein of Kinloch et al. in order to produce a recombinant ZP3

polypeptide or glycopolypeptide. Kinloch et al. provide motivation by teaching the

importance of a specific portion of ZP3 glycopeptide located in the carboxy-terminal end

of both mouse and human ZP3 protein (See p. 267, col. 1, lines 29-51). Kinloch et al.

then sets forth a method of producing mouse and chimeric human ZP3 protein (See p.

264, GLYCOPROTEIN PURIFICATION).

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Claims 71 and 72 are rejected under 35 U.S.C. 103(a) as being unpatentable over Chamberlin & Dean, 87 PROC. NATL. ACAD. Sci. 6014 (1990) in view of Kinloch et al., 92 PROC. NATL. ACAD. Sci. USA 263 (1995) and Rosiere & Wasserman, 154 DEV. BIOL. 309 (1992).

For a discussion of Chamberlin & Dean and Kinloch et al., see supra.

Chamberlin & Dean do not teach a conservatively substituted amino acid sequence of SEQ ID NO:2.

However, Rosiere & Wasserman teach the general location in mouse ZP3 protein sequence responsible for sperm-binding activity (See p. 314, col. 1, lines 31-35). Specifically, Rosiere & Wasserman identify a particular peptide fragment derived from the carboxy-terminal end of the mouse ZP3 protein (See p. 314, col. 2, lines 19-27). It is noted that Applicants' also derived SEQ ID NO: 2 from the carboxy-terminal end of human ZP3 (See specification, p. 6, lines 28-30).

Kinloch et al. extend the knowledge of the art by pointing to the exact amino acid residues of human ZP3 likely responsible for sperm binding (See p. 267, col. 1, lines 29-

51) and describing site-directed ZP3 mutant proteins (See p. 263, col. 2, PLASMID

CONSTRUCTION FOR SITE-DIRECTED MUTAGENESIS) in order to map the mouse ZP3 sperm

binding site.

Therefore, it would have been obvious to a person of ordinary skill in the art to combine

the peptide of Chamberlin & Dean to the ZP3 mutants of Kinloch et al. to provide a

peptide of SEQ ID NO: 2 with conservative amino acid substitutions.

Chamberlin & Dean provide motivation by teaching the similarity between mouse and

human ZP genes (See p. 6014, col. 2, lines 16-18). According to Chamberlin & Dean,

mouse and human ZP3 share 67% identity (See Chamberlin & Dean, p. 6016, col. 2,

lines 14-22). Persons of ordinary skill in the art would recognize this as a high degree

of homology.

Rosiere & Wasserman extend the knowledge in the art by determining the general

location within the mouse ZP3 protein sequence responsible for sperm-binding activity

(See p. 314, col. 1, lines 31-35). Specifically, Rosiere & Wasserman identify a particular

peptide fragment derived from the carboxy-terminal end of the mouse ZP3 protein (See

p. 314, col. 2, lines 19-27). It is noted that Applicants' also derived SEQ ID NO: 2 from

the carboxy-terminal end of human ZP3 (See specification, p. 6, lines 28-30). Rosiere &

Wasserman teach that this fragment from the carboxy-terminal end of ZP3 is heavily

glycosylated and make the following suggestion:

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"Identification and modification of these [glycosylation] sites (e.g., by sitedirected mutagenesis or exon swapping) should lead to a better understanding of the molecular basis of carbohydrate-mediated, speciesspecific gamete interaction in mammals." (p. 316, col. 1, lines 10-14)

Finally, Kinloch et al. extend the knowledge in the art by pointing to the exact amino acid residues of human ZP3 likely responsible for sperm binding (See p. 267, col. 1, lines 29-51), including Ser342 (lines 41-3). When the amino acid sequence of Kinloch et al. (See Fig. 1) is aligned with the sequence of Chamberlin & Dean (See Fig. 3) and the claimed SEQ ID NO:2, it appears that Ser342 aligns with Ser34 of the claimed SEQ ID NO:2.

As evidenced by Kinloch et al., the concept of creating mutant ZP3 proteins and subsequent testing for sperm-binding activity is a well-known and routine practice in the art. Once Rosiere & Wasserman identified the general region on the ZP3 protein likely to be responsible for sperm-binding activity (i.e. the carboxy-terminal region), a person of ordinary skill in the art would have considered the construction of mutant ZP3 proteins, including a conservative substitution at Ser34 of SEQ ID NO:2 as routine experimentation. By Applicants' admission, the choice of amino acid for substitution (i.e. conservative substitutions) is also known in the art (See specification, p. 11, lines 6-20). Therefore, Applicants' claimed polypeptide having conservative amino acid substitutions is an obvious improvement over the teachings of Chamberlin & Dean, Rosiere & Wasserman, and Kinloch et al.

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Claims 71 and 72 recite the claim language "wherein said polypeptide binds human spermatozoa at least ten times as strong as an equivalent molar amount of mouse

ZP3." Examiner interprets this recitation as functional language that is not given

patentable weight.

Response to Arguments

With respect to prior 112, first paragraph rejections, Applicants' arguments filed 10/03/2003 (See "Section V") have been carefully considered and are persuasive in light of subsequent claim cancellations and amendments. With respect to claims 69-72,

the prior 112, first paragraph rejections are withdrawn.

With respect to prior 102(b) and 102(e) rejection based on Dean (US 5,641,487),

Applicants' arguments filed 10/03/2003 (See "Section VI") have been carefully

considered. The prior 102(b) and 102(e) rejection based on Dean (US 5,641,487) is

withdrawn in light of recent amendments to the claims and in light of the rejection

presented in the current action based on Chamberlin & Dean, 87 PROC. NATL. ACAD.

Sci. 6014 (1990). Applicants assert that Dean does not anticipate new claims 69-78

because Dean does not teach the polypeptide of SEQ ID NO:2. Applicants' arguments

have been carefully considered but are not persuasive. Both Dean and Chamberlin &

Dean substantially disclose the claimed SEQ ID NO:2. Applicants are claiming a

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polypeptide comprising SEQ ID NO:2. The use of open claim language "comprising" allows for anticipation by sequences having additional amino acid residues. "When an examiner obtains a product which reasonably appears to fall within the scope of that which is claimed by a patent applicant, it is reasonable to shift the burden to the applicant to provide evidence showing that the product of the prior art does not fall within the scope of applicants' claims." Ex parte Maizel, 27 USPQ2d 1662, 1667 (BPAI 1992) (citing In re Swinehart, 439 F.2d 210, 169 USPQ 226 (CCPA 1971); In re Best, 562 F.2d 1252, 195 USPQ 430 (CCPA 1977) and In re Fitzgerald, 619 F.2d 67, 205 USPQ 594 (CCPA 1980).

With respect to prior 103(a) rejection based on Dean (US 5,641,487) in view of Chamberlin & Dean (87 PRoc. Natl. Acad. Sci. 6014 (1990)), Applicants' arguments filed 10/03/2003 (See "Section VI") have been carefully considered. The prior 103(a) rejection based on Dean (US 5,641,487) in view of Chamberlin & Dean (87 PRoc. Natl. Acad. Sci. 6014 (1990)) is withdrawn in light of amendments to the claims and in light of the rejection presented in the current action based on Chamberlin & Dean (87 Proc. Natl. Acad. Sci. 6014 (1990)) in view of Kinloch et al., 92 Proc. Natl. Acad. Sci. USA 263 (1995) and Rosiere & Wasserman, 154 Dev. Biol. 309 (1992). Applicants have argued that the sperm-binding and acrosome reaction inducing activity of the claimed polypeptide is demonstrated in the specification (See Remarks p. 13, lines 14-15, filed 10/03/2003). Applicants also argue that Chamberlin & Dean do not describe the claimed polypeptides (See Remarks p. 16, lines 7-8, filed 10/03/2003). Applicants'

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arguments have been carefully considered but are not persuasive. A recitation of the intended use of the claimed invention must result in a structural difference between the claimed invention and the prior art in order to patentably distinguish the claimed invention from the prior art. See *In re Casey*, 152 USPQ 235 (CCPA 1967) and *In re Otto*, 136 USPQ 458, 459 (CCPA 1963). Here, as explained *supra*, Applicants' claimed structural differences (i.e. conservative amino acid substitutions) do not distinguish the claimed invention over the prior art because a person or ordinary skill in the art would have been motivated to combine the peptide of Chamberlin & Dean to the ZP3 mutants of Kinloch et al. to provide a peptide of SEQ ID NO: 2 with conservative amino acid substitutions.

#### Conclusion

No claims are allowed.

Applicant's amendment necessitated the new grounds of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the

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shortened statutory period will expire on the date the advisory action is mailed, and any

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extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of

the advisory action. In no event, however, will the statutory period for reply expire later

than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the

examiner should be directed to David J Venci whose telephone number is 571-272-

2879. The examiner can normally be reached on 08:00 - 16:30 (EST). If attempts to

reach the examiner by telephone are unsuccessful, the examiner's supervisor, Long Le

can be reached on 571-272-0823. The fax phone number for the organization where

this application or proceeding is assigned is 703-872-9306.

Information regarding the status of an application may be obtained from the Patent

Application Information Retrieval (PAIR) system. Status information for published

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Center (EBC) at 866-217-9197 (toll-free).

SUPERVISORY PATENT EXAMINER TECHNOLOGY CENTER 1600

David J Venci

Examiner

06/28/01